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(54) ROOM TEMPERATURE SETTING COMPOSITION

(57)Abstract:

PROBLEM TO BE SOLVED: To obtain a room temperature setting composition having a low viscosity, excellent in handleability, mechanical strength and adhesiveness, and enabling the cure rate to be adjusted in a wide range by making the composition include two kinds of specific polyoxyalkylene polymers and an epoxy resin as essential ingredients.

SOLUTION: This composition comprises (A) a polyoxyalkylene polymer having at least one hydrolyzable silicon group of the formula SiX2R1 [R1 is a 1–20C (substituted)monomvalent organic group; X is OH or the like], (B) a polyoxyalkylene polymer having at least one hydrolyzable silicon group of the formula SiX3 which can be obtained by introducing an unsaturated group into the end of a polyoxyalkylene polymer having at least one OH and then reacting with the mercapto group of a silicon compound (e.g. 3– mercaptopropyltrimethoxysilane or the like) of the formula HS-R2-SiX3 (R2 is a 1–17C divalent hydrocarbon) and (C) an epoxy resin (e.g. a bisphenol A-type epoxy resin or the like).

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CLAIMS

[Claim(s)]

unsaturation group and a formula (3) react, and which is expressed with a following formula (2), and a room-temperature-curing nature constituent (D) which uses an epoxy resin (G) as an essential [Claim 1]A polyoxyalkylene polymer (A) which has a hydrolytic silicon group expressed with a following formula (1). And after introducing an unsaturation group into an end of a polyoxyalkylene polymer which has a hydroxyl group, A polyoxyalkylene polymer (B) which has a hydrolytic silicon group which is obtained by making a sulfhydryl group of a silicon compound expressed with this ingredient

- SiX₂R¹ ... (1)

(Among a formula (1), as for substitution of the carbon numbers 1-20 or an unsubstituted univalent organic group, and X, a hydroxyl group or a hydrolytic basis, however two X may be the same, or R^1 may differ.)

- SiX₃ ... (2)

(X are the same as the above among a formula (2).) three X may be the same or may differ.

HS-R²-SiX₃ ... (3)

(X are the same as the above among a formula (3).) three X may be the same or may differ. \mathbb{R}^2 is a [Claim 2] The room-temperature-curing nature constituent (D) containing an epoxy curing agent (E) divalent hydrocarbon group of the carbon numbers 1-17.

[Claim 3]The room-temperature-curing nature constituent (D) containing a compound (F) which has simultaneously a hydrolytic silicon group and reactive functional groups other than hydrolysis nature in the same molecule according to claim 1 or 2. according to claim 1.

[Translation done.]

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DETAILED DESCRIPTION

Detailed Description of the Invention

[Field of the Invention]This invention relates to the room-temperature-curing nature constituent hardened under hygroscopio-surface-moisture existence.

0002]

Description of the Prior Art]The polyoxyalkylene polymer which has a hydrolytic silicon group at the end is used for the use of a coating composition, seal constituents, etc., such as sealant and adhesives, taking advantage of the feature that a hardened material has rubber elasticity. The polymer which has a hydrolytic silicon group which two hydrolytic bases per silicon atom indicated to JP,61–18582,B, JP,3–72527,A, and JP,3–47825,A as such a polymer combine is indicated. Such a polymer is dramatically excellent in the balance of physical properties, such as being able to obtain a flexible hardened material and excelling also in storage stability.

became hyperviscosity even if it originated in the urethane bond itself, it had a problem in workability. silyl groups caused hydrolysis and crosslinking reaction, and a polymer served as hyperviscosity with a small amount of moisture which remains in the system of reaction in that case, plentifully. Since it per silicon atom combine with JP,58-10418,B and JP,58-10430,B --- a molecular weight --- 6000 or less --- the hardenability polymer of low molecular weight is indicated comparatively. Since reactivity quick, but. The molecular weight was inferior to the elongation and pliability of the hardened material sufficient as and is known for promotion of a reaction, was desirable, there was a case where alkoxy [0004]As a method of improving storage stability in the polymer which has a hydrolytic silicon group reaction of polyoxypropylene polyol and an isocyanate substitution type trialkoxysilane compound in [0003]moreover — it is a polymer which has a hydrolytic silicon group which three hydrolytic bases which are acquired from 6000 or less and it being low molecular weight comparatively by hardening. which three hydrolytic bases per silicon atom combine. The polymer obtained by the urethane-ized etc. by one side in the polymer which has a hydrolytic silicon group which two hydrolytic bases per JP,10-245482,A, The constituent which consists of a curing catalyst and amino group substitution However, although a cure rate can be made to some extent quick by selection of a curing catalyst groups are an alkyl dialkoxy group is indicated. The polymer intrinsically obtained by the urethaneisocyanate group in this method is low and to obtain significant reaction velocity, Although use of reaction accelerators, such as various kinds of metal salt which a urethane-ized reaction may be silicon atom combine, Making it improve by leaps and bounds had a limit from reactivity with the hydrolytic bases per such a silicon atom combine has the characteristic that a cure rate is very polyoxyalkylene polymer and molecular terminal whose molecular terminal is the Tori alkoxy silyl compound in this method is used. In order the reactivity of the polyoxypropylene polyol and the water of a silicon group, and the limit was among the uses asked for a quick cure rate naturally. with water is high, the hardenability polymer which has a hydrolytic silicon group which three ized reaction of polyoxypropylene polyol and an isocyanate substitution type trialkoxysilane From performance in use being high, industrial production was actually carried out and the commercial scene is established in the use of adhesives, a water blocking material, etc. alkoxysitane, Or in JP,10-245484,A, the mixture of the polyoxyalkylene polymer whose

[0005]By one side, it could not say that the polyoxyalkylene polymer which has a hydrolytic silicon group at the above-mentioned end was never excellent about the intensity of a hardened material,

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JP,2001-072855,A [DETAILED DESCRIPTION]

and adhesive strength with adherend, but the use was restricted naturally. As a method of improving an adhesive property, the method of using together the polymer which has a hydrolytic silicon group, and an epoxy resin is proposed by JP,61-268720,A and JP,7-2828,B. Although it was effective in such a method improving the intensity and adhesive strength of a hardened material, in respect of the cure rate, it was not yet enough, and when using it especially as adhesives, the adhesive property was wanted to be revealed in shortest possible time.

[0006]And it applies for the room-temperature-curing nature constituent which uses as an essential ingredient the polymer and epoxy resin which have a hydrolytic silicon group which three hydrolytic bases per silicon atom combine in ten to Japanese-Patent-Application-No. 204041 specification by these people. As a polyoxyalkylene polymer which has a hydrolytic silicon group which three hydrolytic bases per silicon atom combine in this method, Although the polymer obtained by making the trimethoxysilane which is a hydrosilyl compound react to the end of a raw material polyoxyalkylene polymer to the polymer which introduced the unsaturation group is used, The trimethoxysilane which is the hydrosilyl compound used here had big difficulty on [, such as a transfer except of storage,] handling art from the problem of safe according to the disproportionation to Silang

[0007]In this method, when hydrolysis and crosslinking reaction of a hydrolytic silicon group advanced promptly, while the oure rate of resin became quick, available time and what is called an open time had the fault. I that it is short I of having elapsed and being hard to use it. Adjustment of adjusting freely generally, so that adjustment of the oure rate did not receive influence in the reactivity of the end although the thing possible to some extent was known by the kind of the various metal carboxylate which has a catalysis of a hardening reaction, basicity, an acidic compound, etc., or selection of quantity it is called for that it can reappord to various cure rates also industrially, and controllable art was freely searched for about hardenability.

Uponon (s) to be Solved by the Invention]In view of the above-mentioned problem, the polymer which has a hydrolytic silicon group industrially obtained easily considering the silane compound [handling / a silane compound] as a raw material is used for the purpose of this invention, It is dealt with by hypoviscosity, and excels in a sex, and excels in the mechanical strength of the hardened material after hardening, and adhesive strength with adherend, and is in moreover providing the hardeneability constituent which can be adjusted in the range with a wide cure rate.

by the standard of the problem]Namely, a polyoxyalkylene polymer (A) in which this invention has a hydrolytic silicon group expressed with a following formula (1). And after introducing an unsaturation group into an end of a polyoxyalkylene polymer which has a hydroxyl group. Are obtained by making a sulfhydryl group of a silicon compound expressed with this unsaturation group and a formula (3) react. A polyoxyalkylene polymer (B) which has a hydrolytic silicon group expressed with a following formula (2), and a room-temperature-curing nature constituent (D) which uses an epoxy resin (C) as an essential ingredient are provided.

[0010]- SiX2R1 ... (1)

(Among a formula (1), as for substitution of the carbon numbers 1–20 or an unsubstituted univalent organic group, and X, a hydroxyl group or a hydrolytic basis, however two X may be the same, or \mathbb{R}^1 may differ.)

- SiX, ..

(X are the same as the above among a formula (2).) three X may be the same or may differ.

(X are the same as the above among a formula (3).) three X may be the same or may differ. \mathbb{R}^2 is a

HS-R²-SiX₃ ... (3)

divalent hydrocarbon group of the carbon numbers 1–17. [0011] [0011] [Embodiment of the Invention](Polyoxyalkylene polymer) The polyoxyalkylene polymer (B) which has a hydrolytic silicon group expressed with the polyoxyalkylene polymer (A) and the above-mentioned formula (2) which have a hydrolytic silicon group expressed with the above-mentioned formula (1) used in this invention so that it may state below, It is preferred to use as a raw material the

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polyoxyalkylene polymer which has a functional group, to introduce a hydrolytic silyl group into a part or all of the functional group via an organic group, and to be manufactured, For example, what is indicated to JP,3-47825,A, JP,3-72527,A, and JP,3-79627,A is mentioned.

(Raw material polyoxyalkylene polymer) As the polyoxyalkylene polymer (A) used in this invention, and a raw material polyoxyalkylene polymer of (B), the thing of the hydroxyl group end which makes cyclic ether etc. react and is manufactured is preferred under existence of a catalyst and existence of an initiator. This raw material polyoxyalkylene polymer may be on a straight chain, may be a letter of branching, or there may be with these mixtures.

[0012]As an initiator, the hydroxy compound etc. which have one or more hydroxyl groups can be used. As cyclic ether, ethylene oxide, propylene oxide, butylene oxide, hexyleneoxide, a tetrahydrofuran, etc. are mentioned. As a catalyst, alkaline metal catalysts, such as a potassium system compound and a caesium system compound, a composite metal cyanide complex catalyst, a metalloporphyrin catalyst, etc. are mentioned.

[0013]As a functional group number, the raw material polyoxyalkylene polymer of 2-8 is preferred, and when a functional group number uses the thing of 2 or 3 as a raw material, it is especially desirable from the polyoxyalkylene polymer (A) and (B) which demonstrates the adhesive strength which was excellent greatly in pliability as the hardened material characteristic being obtained, as a raw material polyoxyalkylene polymer, the thing of the amount of polymers is preferred — further the ratio of weight average molecular weight (Mw) and a number average molecular weight (Mn) — the narrow thing of molecular weight distribution with small Mw/Mn is preferred.

the narrow thing of molecular weight distribution with small Mw/Mn is preferred.

[0014]When the raw material polyoxyalkylene polymer of Mw/Mn which is different with the same number average molecular weight (Mn) compares, it has the feature that the viscosity of a polymer becomes low and the thing which has small Mw/Mn is excellent in workability. When it is going to adjust the viscosity of a polymer to a certain fixed level, the content of the polymer of low molecular weight decreases as the small thing of Mw/Mn. The polyoxyalkylene polymer which has by this a hydrolytic silicon group obtained considering this as a raw material. As for the hardened material produced by excelling in the hardening characteristics — the depths hardenability at the time of hardening becomes good — compared with the case where the large thing of Mw/Mn is used as a raw material, and hardening, the elongation of a hardened material serves as high intensity large also what has a the same elastic modulus.

Incornage material polyoxyalkylene polymer of the amount of polymers, it can obtain by the method of using a composite metal cyanide complex catalyst, the method of quantifying many and carrying out polymers quantification by [which manufactured using the alkali catalyst etc.] making many halogenated compounds, such as a methylene chloride, the polyoxyalkylene polymer of low molecular weight react comparatively, etc. As for the small raw material polyoxyalkylene polymer of Mw/Mn, what is obtained considering a composite metal cyanide complex as a catalyst is preferred. The complex which uses zinchexacyano obaltate as the main ingredients as a composite metal cyanide complex is preferred, and ether and/or an alcoholic complex are especially preferred. The presentation can use what is intrinsically indicated to JP.46-27250,B. In this case, as ether, ethyleneglycol dimethyl ether (glyme), diethylene glycol dimethyl ether (glyme), diethylene glycol dimethyl ether (glyme), and especially glyme is preferred from a point of the handling at the time of manufacture of a

complex. As aloohol, t-butanol is preferred. [0016]As a molecular weight of a raw material polyoxyalkylene polymer, the thing of 6,000–50,000 is preferred at a number average molecular weight (Mn), and, specifically, especially the thing of 8,000–20,000 is preferred. As Mw/Mn of a raw material polyoxyalkylene polymer, 1.7 or less are preferred, 1.6 or less are still more preferred, and 1.5 especially or less are preferred.

[0017]As a raw material polyoxyalkylene polymer, the copolymer of a polyoxyethylene, polyoxybethylene, polyoxy butylene, polyoxy hexylene, polyoxy tetramethylen, and two or more sorts of cyclic ether is specifically mentioned. Especially a desirable raw material polyoxyalkylene polymer is the polyoxypropylene polyol of 2 – 6 value, and are polyoxypropylene diol and polyoxypropylene triol especially. When using for the method of the following (b) or (**k), the polyoxyalkylene polymer of unsaturation groups, such as an allyl end polyoxypropylene monocar, can also be used. (Polyoxyalkylene polymer (A) has a hydrolytic silicon group expressed with the end or side chain of a chain with a following formula (1).

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JP,2001-072855,A [DETAILED DESCRIPTION]

(Among a formula (1), as for the substitution of the carbon numbers 1–20 or an unsubstituted univalent organic group, and X, a hydroxyl group or a hydrolytic basis, however two X may be the same, or \mathbb{R}^1 may differ.)

R^{jin} formula (1) is a univalent organic group the substitution of the carbon numbers 1–20, or unsubstituted, and a with a carbon number of eight or less alkyl group, a phenyl group, or a fluoro alkyl group is preferred. A methyl group, an ethyl group, a propyl group, a butyl group, a hexyl group, a cyclohexyl group, a perifically, especially a methyl group is

preferred, when two or more R¹ exist, those R¹ may be the same, or may differ.

[0019]As a hydrolytic basis in X, a halogen atom, an alkoxy group, an acyloxy group, an alkenyloxy group, a carbamoyl group, an amino group, an aminooxy group, a KETOKISHI mate group, etc. are mentioned, for example. As for the carbon number of the hydrolytic basis which has a carbon atom among these, six or less are preferred, and four especially or less are preferred. The viewpoint of hydrolysis nature being quiet and being easy to deal with it among these to especially an alkoxy group is preferred. Especially as an alkoxy group, a methoxy group and an ethoxy basis are preferred, and a methoxy group is the most preferred.

[0020]That is, it is preferred that it is especially the structure of having an alkyl dialkoxy silyl group, as a hydrolytic silicon group expressed with a following formula (1), and a methyl dimethoxy silyl group is the most preferred. The hydrolytic silicon group expressed with a formula (1) is usually introduced into a raw material polyoxyalkylene polymer via an organic group. That is, as for a polyoxyalkylene polymer (A), it is preferred to have a basis expressed with a formula (4).

[0021]-R⁰-SiX₉R¹ ... (4)

(A divalent organic group, R^1 , and X of R^0 are the same as that of the above among a formula (4).) Although the method in particular of introducing a hydrolytic silicon group to a raw material polyoxylatylene polymer is not limited, it can be introduced, for example by (b) of the following – the mathod of $\{x*\}$

(**) A method to which the hydrosilyl compound expressed with it by a formula (5) after introducing an unsaturation group into the end of the polyoxyalkylene polymer which has a hydroxyl group is made to react.

[0022]HSiX,R¹ ... (5)

(R¹ and X are the same as the above among a formula (5).)

The method of making the compound which has an unsaturation group and a functional group react to the terminal hydroxyl groups of the polyoxyalkylene polymer which has a hydroxyl group, and combining it by the ether bond, the ester bond, a urethane bond, or carbonate combination as a method of introducing an unsaturation group, is mentioned. As an unsaturation group here, CH₂=CH-

R'- (R' is a divalent hydrocarbon group of the carbon numbers 1–18) is preferred. As for R', it is preferred that it is a hydrocarbon group of the carbon numbers 1–5. Especially as an unsaturation group, an allyl group is preferred. Especially as a compound which has an unsaturation group and a functional group, allylchloride is preferred, and the hydroxyl group of the polyoxyalkylene polymer which has a hydroxyl group can be used as an allyloxy group by using this. When polymerizing alkylene oxide, the method of introducing an unsaturation group into the side chain of a raw material polyoxyalkylene polymer can also be used by adding and carrying out copolymerization of the unsaturation group content epoxy oompounds, such as allyl glycidyl ether.

[0023]When making a hydrosilyl compound react, catalysts, such as a platinum system catalyst, a rhodium system catalyst, a cobalt system catalyst, a palladium system catalyst, and a nickel series catalyst, can be used. Platinum system catalysts, such as chloroplatinic acid, platinum metal, a platinum chloride, and a platinum olefin complex, are preferred. As for the reaction to which a hydrosilyl compound is made to react, it is preferably preferred to carry out at the temperature of 60-120 ** for several hours 30-150 **.

[0024](**) A method to which the compound expressed with the end of the polyoxyalkylene polymer which has a hydroxyl group by a formula (6) is made to react.

R¹-SiX₂-R²NCO ... (6)

(R $^{\rm l}$ and X are the same as the above among a formula (6).) ${
m R}^2$ is a divalent hydrocarbon group of the

JP,2001-072855,A [DETAILED DESCRIPTION]

carbon numbers 1-17.

4 publicly known urethane-ized catalyst may be used in the case of the above-mentioned reaction. As for the above-mentioned reaction, it is preferably preferred to carry out at the temperature of 50-150 ** for several hours 20-200 **.

of the polyoxyalkylene polymer which has a hydroxyl group and considering it as an isocyanate group end, Method R¹-SiX,-R²W to which W basis of the silicon compound expressed with a formula (7) to [0025](**) After making polyisocyanate compounds, such as tolylene diisocyanate, react to the end this isocyanate group is made to react ... (7)

 $(R^1, R^2,$ and X are the same as the above among a formula (7).) Active hydrogen containing group as which W is chosen from a hydroxyl group, a carboxyl group, a sulfhydryl group, and an amino group (the 1st class or the 2nd class).

catalyst, etc. of a peroxide system, azo, or a redox system are mentioned, for example. Specifically as radiation or heat in the case of the above-mentioned reaction, without using a polymerization initiator Polymerization initiators, such as a radical generator, may be used, and it may be made to react with that of what was explained in (b). 3-mercapto propylmethyl dimethoxysilane etc. are mentioned as a peroxide, t-alkyl peroxy ester, acetyl peroxide, diisopropyl peroxy carbonate, etc. are mentioned. As expressed with the formula (7) whose W is a sulfhydryl group are made to react after introducing an [0026]The method and unsaturation group which introduce an unsaturation group are the same as for the above–mentioned reaction, it is preferably preferred to carry out at 50–150 ** for several depending on the case. As a polymerization initiator, a polymerization initiator, a metal compound (**) A method to which the unsaturation group and the sulfhydryl group of a silicon compound a polymerization initiator, 2,2"-azobisisobutyronitrile, 2,2"-azobis 2-methylbutyronitrile, benzoyl unsaturation group into the end of the polyoxyalkylene polymer which has a hydroxyl group. silicon compound by which W is expressed with the formula (7) which is a sulfhydryl group. hours - tens hours 20-200 **.

(Polyoxyalkylene polymer (B)) A polyoxyalkylene polymer (B) has a hydrolytic silicon group expressed with the end or side chain of a chain with a following formula (2).

[0027]- SIX₃ ... (2)

mentioned formula (2) has the characteristic that the cure rate at the time of hardening is very quick. alkoxy group, an acyloxy group, An alkenyloxy group, a carbamcyl group, an amino group, an aminooxy group, a KETOKISHI mate group, a hydride group, etc. are mentioned, an alkoxy group is preferred in these, a methoxy group and an ethoxy basis are specifically preferred, and a methoxy group is the (X are the same as the above among a formula (2).) three X may be the same or may uniter. Since reactivity is high, the polymer which has a hydrolytic silicon group expressed with the above— As a hydrolytic basis in X in a formula (2), As well as the above, for example, a halogen atom, an are the same as the above among a formula (2).) three X may be the same or may differ. most preferred.

most preferred. As for a polymer (B), it is preferred that the hydrolytic silicon group expressed with a [0028]That is, it is preferred that it is especially the structure of having the Tori alkoxy silyl groups, polymer (A). That is, as for a polyoxyalkylene polymer (B), it is preferred to have a basis expressed as a hydrolytic silicon group expressed with a following formula (2). A trimethoxysilyl group is the formula (2) is introduced into a raw material polyoxyalkylene polymer via an organic group like a with a formula (8).

[0029]-Y-S-R²-SiX₃ ... (8)

divalent hydrocarbon group of 1-17. R² is a divalent hydrocarbon group of the carbon numbers 1-17. As Y, the hydrocarbon group of the carbon numbers 1-5 is preferred, and especially a trimethylene (X are the same as the above among a formula (8).) three X may be the same or may differ. Y is a

[0030]A polyoxyalkylene polymer (B) is obtained by making the sulfhydryl group of a silicon compound expressed with this unsaturation group and a formula (3) react, after introducing an unsaturation group into the end of the polyoxyalkylene polymer which has a hydroxyl group. group is preferred. As R^2 , the hydrocarbon group of the carbon numbers 1–5 is preferred, and especially a trimethylene group is preferred.

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(X are the same as the above among a formula (3).) three X may be the same or may differ. \mathbb{R}^2 is a divalent hydrocarbon group of the carbon numbers 1-17.

About the details of the introducing method, it is the same with having explained in method (**) which introduces the hydrolytic silicon group of said polyoxyalkylene polymer (A). As an unsaturation group, CH,=CH-R'- (R' is the same as the above) is preferred as above-mentioned.

Novolak type epoxy resin, such as bisphenol F novolak type epoxy resin, A hydrogenation bisphenol A reactivity when hardening, and a hardened material is especially preferred from points -- it is easy to mercaptopropyl triethoxysilane, etc. are mentioned. It has the characteristic that the cure rate of the polyoxyalkylene polymer obtained with a described method is large, and has the feature of excelling in resin, a resorcinol type epoxy resin, and petroleum resin, etc. are raised, It is not limited to these. A bisphenol A type epoxy resin, bisphenol F type epoxy resin, novolak type epoxy resin, and especially as tetrahydrophtal acid diglycidyl and hexahydrophthalic acid diglycidyl, m~aminophenol series epoxy type epoxy resin, the glycidyl ether type epoxy resin of bisphenol A / propylene oxide addition, 4-glycidyloxy benzoic acid glycidyl, phthalic acid diglycidyl. Diglycidyl ester system epoxy resins, such (Epoxy resin (C)) As an apoxy resin (C) used for this invention, A publicly known thing can be used [0031]As a silicon compound expressed with a formula (3), 3-mercapto propyltrimethoxysilane, 3widely conventionally and it is in a concrete target. Bisphenol A type epoxy resin, Fire retardancy type epoxy resins, such as bisphenol F type epoxy resin and a glycidyl ether type epoxy resin of triglycidyl isocyanurate, polyalkylene glycol diglycidyl ether, and glycerin, a hydantoin type epoxy epoxidation thing of unsaturation polymers, such as glycidyl ether of polyhydric alcohol, such as resin, a diaminodiphenylmethane system epoxy resin, Glycidyl ester-typed-epoxy-resin, glycidyl amine-type-epoxy-resin, isocyanurate type epoxy resin, urethane modified epoxy resin, various tetrabromobisphenol A, Gresol novolak type epoxy resin, bisphenol A novolak type epoxy resin, cycloaliphatic-epoxy-resin, and N.N-diglycidyl aniline, N.N-diglycidyl o-toluidine, Although the [0032]As an epoxy resin (C), what contains two or more epoxy groups in a molecule has high workability by hypoviscosity. Acquisition of a raw material is also easy and industrially useful. the epoxy resin of a phthalic acid diglycidyl ester system are [among these] preferred. build the three-dimensional network structure.

(Epoxy curing agent (E)) As for the room-temperature-curing nature constituent (D) of this invention. it is still more preferred to contain an epoxy curing agent (E). As an epoxy curing agent (E),

system polymers (end amination polyoxypropylene glycol.) which average the basis which can react to an epoxy group and it has at least one piece in intramolecular, such as phenoxy resin, carboxylic acid, group, Although liquefied end functional group content polymers etc. which were embellished with the Triethylenetetramine, tetraethylenepentamine, diethylamino propylamine, N-aminoethyl piperazine, misophoronediamine, Amines or those salts, such as 2,4,6-tris(dimethyl aminomethyl) phenol, Aldimine, compounds, phthalic anhydride, a hexahydrophthalic anhydride, A tetrahydrophthalic anhydride and and alcohols An end end carboxylation polyoxypropylene glycol etc. A hydroxyl group, a carboxyl amino group eto., such as polybutadiene, hydrogenation polybutadiene, an acrylonitrile butadiene enamines, polyamide resin, imidazole derivatives, and dicyandiamides. Boron trifluoride complex succinic anhydride, pyromellitic dianhydride, and anhydrous KUROREN acid, polyalkylene oxide methylene tetrahydro phthalic anhydride, Anhydrous carboxylio acid, such as a DODESHINIRU copolymer, and an acrylic polymer, are raised, it is not limited to these. 2,4,6-tris(dimethyl xylylene diamine, m-phenylenediamine, Diaminodiphenylmethane, diaminodiphenyl sulfone, conventionally, can use a publicly known thing widely and specifically Diethylenetriamine, aminomethy!) phenol is [among these] the most preferred.

[0033]A compound like ketimines as shown with the following formula (9) or (10), and silazanes can also be used.

R3R4C=N-R5-NH-R6-N=CR7R8 ... (9)

(The inside of a formula, $\rm R^3$, $\rm R^4$, $\rm R^7$, and $\rm R^8$ are a hydrogen atom, a halogen atom, or a univalent hydrocarbon group.) R⁵ and R⁶ are divalent hydrocarbon groups.

R9Rf0C=N-R11-N=CR12R13 ... (10)

(The inside of a formula, $m R^9$, $m R^{10}$, $m R^{12}$, and $m R^{13}$ are a hydrogen atom, a halogen atom, or a univalent hydrocarbon group.) \mathbb{R}^{11} is a divalent hydrocarbon group.

Since amine was generated only after these compounds reacted to moisture, when it can use it as an insidious hardening agent and is especially used in combination of one component type, etc., as compared with other epoxy curing agents, it has the feature that storage stability is excellent. [0034]Although the compound etc. which are obtained by the dehydration of carbonyl compounds, such as polyamine, methyl ethyl ketone, etc., such as diethylenetriamine, are raised as a typical example, it is not limited to this. The amino group of the ketimines shown by the above—mentioned formula (9) Styrene oxide, Butyl glycidyl ether, 3-glycidyloxypropyl trinethoxysilane. The compound made to react to epoxy group containing compounds, such as 3-glycidyloxy propylmethyl dimethoxysilane and 3-glycidyloxy propyl triethoxysilane, and mono-isocyanate compounds, such as a phenylisocyanate, can also be used.

(Compound (F) which has simultaneously a hydrolytic silicon group and a reactive functional group besides hydrolytic silicon Motomochi) The room-temperature-curing nature constituent (D) of this invention may contain further the compound (F) which has simultaneously a hydrolytic silicon group and a reactive functional group besides hydrolytic silicon Motomochi. A compound (F) is a compound which has a different functional group of at least two kinds of reactivity in one molecule, and makes a hydrolytic silyl group indispensable, and means the compound usually known as a silane coupling agent or an adhesion grant agent. As a compound (F), a conventionally publicly known compound which is indicated, for example to "the optimal selection and use art of a coupling agent, and an appraisal method (TECHNICAL INFORMATION INSTITUTE)" etc. can be used. When a room-temperature-curing nature constituent (D) contains a compound (F), it is desirable from adhesive strength with adherend acting as Kougami further.

[0035]As a hydrolytic silicon group in a compound (F), the hydrolytic silicon group expressed with a formula (1) and a formula (2) is preferred. As a reactive functional group besides hydrolytic silicon Motomochi in a compound (F), an epoxy group, an amino group, a sulfhydryl group, an acryloxy (meta) group, a carboxyl group, etc. are preferred.

[0036]As a compound (F), epoxy group content Silang, amino group content Silang, sulfhydryl group content Silang, acryloxy (meta) group content Silang, and carboxyl group content Silang are mentioned. As epoxy group content Silang, specifically 3-glycidyloxypropyl trimethoxysilane, 3-glycidyloxy propylmethyl dimethoxysilane, 3-glycidyloxy propyl triethoxysilane, 3-(N,N-diglycidyl) aminopropyl trimethoxysilane, N-glycidyl N,N-bis[3-(methyl dimethoxy silyl) propyl] amine, N-glycidyl N,N-bis[3-(trimethoxysilyl) propyl] amine, etc. are mentioned.

N.N-bis[3-(trimethoxysilyl) propyl] amine, etc. are mentioned. [0037]As amino group content Silang, specifically 3-aminopropyl trimethoxysilane, 3-aminopropyl triethoxysilane, 3-aminopropyl triethoxysilane, 3-aminopropyl methyl dimethoxysilane, N-(2-aminopropyl trimethoxysilane, N-(2-aminoethyl)-3-aminopropyl trimethoxysilane, N-(2-aminoethyl)-3-aminopropyl triethoxysilane, N,N-bis[3-(methyl dimethoxy silyl) propyl] amine, N,N-bis[3-(methyl dimethoxy silyl) propyl] ethylenediamine, N,N-bis[3-(trimethoxysilyl) propyl] ethylenediamine, N,N-bis[3-(trimethoxysilyl) propyl] diethylenetriamine $[H_2N(C_2H_4NH)_2C_3H_6Si(OCH_3)_3]$. N-[(3-trimethoxysilyl) propyl] triethylenetetramine $[H_2N(C_2H_4NH)_3C_3H_6Si(OCH_3)_3]$. 3-ureido propyl triethoxysilane, N-(N-vinylbenzyl 2-aminoethyl)-3-aminopropyl trimethoxysilane, 3-anilino propyltrimethoxysilane etc. are

vinylbenzyl 2-aminoethyl)-3-aminopropyl trimethoxysilane, 3-anilino propyltrimethoxysilane etc. are mentioned.
[0038]As sulfhydryl group content Silang, 3-mercapto propyltrimethoxysilane, 3-mercaptopropyl triethoxysilane, 3-mercapto propylmethyl dimethoxysilane, 3-mercapto propylmethyl diethoxy SHISHIRAN, etc. are specifically raised. As (meth)acryloyloxy group content Silang, 3-methacryloyl

oxypropyl trimethoxysilane, 3-acryloyloxypropyl trimethoxysilane, 3-methacryloyl oxypropyl methyl

dimethoxysilane, etc. are specifically mentioned. [0039]As carboxylane, etc. are specifically mentioned. [0039]As carboxyl group content Silang, specifically 2-carboxyethyl triethoxysilane, 2-carboxyethyl phenylbis(2-methoxyethoxy)Silang, N-(N-carboxylmethyl 2-aminoethyl)-3-aminopropyl trimethoxysilane, etc. are mentioned. The reactant produced by making two or more sorts of these compounds react may be used. As an example of a reactant, the reactant of amino group content Silang and epoxy group content Silang. The reactant of epoxy group content Silang the reactant of amino group content Silang and (meth)acryloyloxy group content Silang, etc. are mentioned. These reactants are easily obtained by mixing the above-mentioned compound and agitating in a room temperature -150 ** temperature requirement for 1 to 8 hours.

[0040]A compound (F) may be used alone and may be used together two or more kinds.

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Room-temperature-curing nature constituent (D)) The polyoxyalkylene polymer (A) in the room-

(Room-temperature-curing nature constituent (D)) The polyoxyalkylene polymer (A) in the room-temperature-curing nature constituent (D) of this invention and the rate of (B) can be arbitrarily chosen according to a use, the characteristic to need, etc.

hardenability is quick, the effect that the manifestation of the adhesive intensity after construction is be moved, and a short time until an adhesive property is revealed especially. On the other hand, from the thing which has a large rate of a polyoxyalkylene polymer (A) being excellent about the extension 0041]Although the polyoxyalkylene polymer (A) and the rate of (B) are arbitrary, (A):(B) =5 - 95:95 arbitrary to the thing conventionally near equivalent hardenability can be obtained. When what has a 70:70 – 30 are the most preferred. That is, adjustment of hardenability becomes possible in the wide characteristic of a hardened material. It is important to obtain the optimal constituent each time by etc., it is useful to paste up in the directions for use which must be fixed so that adherend may not 5 are preferred at a weight ratio, (A):(B) =20 - 80.80 - 20 are still more preferred, and (A):(B) =30 - 20quick is acquired, In the case where he would like to make hardening under low temperature quick range by adjusting a polyoxyalkylene polymer (A) and the rate of (B) arbitrarily. Hardenability can lessening the rate of a polyoxyalkylene polymer (B), and the constituent which has hardenability changing a polyoxyalkylene polymer (A) and the rate of (B) arbitrarily according to a use or the large rate of a polyoxyalkylene polymer (B) is used as adhesives, a sealing material, etc., Since specifically be made quick, so that the rate of a polyoxyalkylene polymer (B) is enlarged, it is characteristic to need.

[0042]Although the rate of the polyoxyalkylene polymer (A) and the epoxy resin (C) to (B) in the room-temperature-curing nature constituent (D) of this invention can be chosen arbitrarily, 1 – 300 weight section is preferred to a polyoxyalkylene polymer (A) and a total of 100 weight sections of (B), and especially 1 – 100 weight section is preferred. When less than this, the intensity of the hardened material after hardening becomes insufficient, and an epoxy resin (C) becomes insufficient [duotility] in being more than this, and is not preferred.

[0043]Although the rate of the polyoxyalkylene polymer (A) and the compound (F) to (B) in the room-temperature-curing nature constituent (D) of this invention can be chosen arbitrarily, 0 – 30 weight section is preferred to a total of 100 copies of a polyoxyalkylene polymer (A) and (B), and especially 0.1 – 10 weight section is preferred. When a compound (F) is added across a mentioned range, it is not desirable from properties balance, such as the handling nature of a constituent, being spoiled, and the cost of a constituent becoming high etc.

[0044]Athough it is not indispensable, the room-temperature-curing nature constituent (D) of this invention does not need to be included even if the additive agent as shown below is included. Hereafter, an additive agent is explained.

Hereafter, an additive agent is explained. (Bulking agent can be used as a bulking agent. Especially the amount (Bulking agent) A publicly known bulking agent can be used as a bulking agent used has 50 – 250 preferred weight section 0.001 to 1000 weight section to a polyoxyalkylene polymer (A) and a total of 100 weight sections of (B). The following are mentioned as an example of a bulking agent. These bulking agents may be used independently and may be used together two or more sorts.

[0045]Heavy calcium carbonate with a mean particle diameter of 1–20 micrometers, precipitated calcium carbonate with a mean particle diameter of 1–3 micrometers manufactured with the sedimentation method. The colloid calcium carbonate which carried out the surface treatment of the surface with fatty acid or a resin acid system organic matter, Calcium carbonate, such as minor nature calcium carbonate, furnes silica, sedimentation nature silica, Surface siliconization silica pulverized coal, a silicic acid anhydride, hydrous silicic acids, and carbon black, Magnesium carbonate, diatomite, calcination or lay, clay, talc, titanium oxide, bentonite, bentonite, ferric oxide, a zinc oxide, an active white, resin beads, wood flour, pulp, a cotton chip, mica, and the blacking wash farina— tubbing — powder state bulking agents, such as farina, graphite, aluminum impalpable powder, and the Flint powder. Fibrous fillers, such as asbestos, glass fiber, a glass filament, carbon fiber, the Kevlar textiles, and a polyethylene fiber.

[0046]The publicly known hollow body of minerals or the quality of organicity can be used. Since specific gravity is low, workability of a hollow body improves — a weight saving and the cobwebbing nature of a constituent are improved in a constituent and its hardened material. As an inorganic hollow body, can illustrate a silicio acid system hollow body and a non-silicio acid system hollow body and as a silicio acid system hollow body. A milt balloon, perlite, glass balloons, a silica balloon, and fly

carbon balloon, etc. can be illustrated, a mit balloon and especially glass balloons are preferred, and balloons, As a non-silicic acid system hollow body, an alumina balloon, a zirconia balloon, a glass balloons are the most preferred.

average particle density, The mean particle diameter of about 10-500 micrometers preferably Usually, [0047]Although there is no limitation in particular in the mean particle diameter of glass balloons, and about 30–100 micrometers, The average particle density of about 0.1– $0.6 \mathrm{g/cc}$ preferably About 0.15– 0.3g/cc. about 0.05-0.5g/cc ***** -- desirable -- about 0.07-0.3g [cc] /and a pressure

glass balloons, etc., the thing nearer to a real ball has the more preferred viscosity of a constituent esistance [of 10-1000kg/cm] ² grade — a 15 - 300 kg/cm² grade and the thing of not less than 90% of ***** are preferably common. Although there is no limitation in particular in the shape of from becoming low.

is raised, and it may be made to foam, after blending what that to which it foamed beforehand may be thermosetting resin, and the hollow body of thermoplastics which constructed the bridge can also be used. As particles, the porous empty capsid which has a hole what is called other than a hollow body balloon, an epoxy balloon, and a urea balloon as a hollow body of thermoplastics, A saran balloon, a [0048]As a hollow body of the quality of organicity, the hollow body of thermosetting resin and the hollow body of thermoplastics can be illustrated, As a hollow body of thermosetting resin, a phenol polystyrene balloon, a polymethacrylate balloon, a polyvinyl alcohol balloon, and a styrene acrylic balloon can be illustrated. What coated the surface of the hollow body of thermoplastics with sufficient as, and contains a foaming agent

[0049]When the hollow body of the quality of organicity is used, the hardenability constituent which ingredients with granular materials, such as calcium carbonate, talc, and titanium oxide, is preferred constituent and its hardened material can be obtained. It is raised by the hollow body of the hybrid has the feature of excelling in elongation with a low modulus in the tractive characteristics of the specifically. What coated the surface of the hollow body which uses polyacrylonitrile as the main hardened material after hardening besides the effect which carries out the weight saving of a type which coated the surface of the organic hollow body with inactive inorganic powder, and from the good thing of familiarity by other bulking agents.

GLASS MODULES (PITTSBURGH CORNING CORP.) etc., As fly ash balloons, CEROSPHERES (PFA MARKETING LTD.), FILLITE (FILLITE U.S.A. INC.) etc., As an alumina balloon, as BW (Showa Denko) and a zirconia balloon, HOLLOW ZIRCONIUMSPHERES (ZIRCOA) etc., As a carbon balloon, [0050]As an example of an inorganic hollow body, as a milt balloon, for example a win light (LUICHI Chemicals) etc., As glass balloons, it is a Scotch whisky light. Glass BABURUZU (3M), CEL-STAR (Tokai industry), Q-CEL (Pacific chemicals), MICRO BALLOON (EMERSON & CUMING), CELAMIC KUREKASU fair (Kureha chemicals), car boss fair (GENERAL TECHNOLOGIESCORP.), etc. are

[0051]As an example of the hollow body of the quality of organicity, as a phenol balloon, for example PHENOLIC MICROBALLOONS (UCC) etc., As an epoxy balloon, ECCOSPHERES EP (EMERSON & CUMING) etc., As a urea balloon, ECCOSPHERES VF-0 (EMERSON & CUMING) etc., As a saran Matsumoto microsphere (Matsumoto Yushi-Seiyaku), etc., As a polystyrene balloon, DYLITE EXPANDABLE POLYSTYRENE (ARCO POLYMERS INC.), EXPANDABLE POLYSTYRENE BEADS balloon, SARANMICROSPHERES (DOWCHEMICAL, COMPANY), Expancel (Japanese phyllite), the (BASF WYANDOTE CORP.) etc. are raised for SX863 (P) (Japan Synthetic Rubber) etc. as a constructed type styrene acrylic acid balloon of a bridge.

inactive inorganic powder, Matsumoto microsphere MFL series (Matsumoto Yushi-Seiyaku) etc. can [0052]As a hollow body of the hybrid type which coated the surface of the organic hollow body with be illustrated, 0.3 - 40 weight section is still more preferably preferred [the amount of the hollow body used] 0.1 to 50 weight section preferably 0.01 to 100 weight section to a polyoxyalkylene

polymer (A) and a total of 100 weight sections of (B). [0053]If a point with required taking care that a hollow body does not break according to the shearing (Plasticizer) A publicly known plasticizer can be used as a plasticizer. The amount of the plasticizer used has 0.001 -- 1000 preferred weight section to a polyoxyalkylene polymer (A) and a total of 100 force at the time of mixing in mixing of a hollow body, especially the hollow body of comparatively weak construction material like glass balloons is removed, it can treat like the usual bulking agent. weight sections of (B). The following are mentioned as an example of a plasticizer. http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fwww4.ipdli... 2010/03/03

JP,2001-072855,A [DETAILED DESCRIPTION]

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[0054]Phthalio ester, such as di∹(2–ethylhexyl)phthalate, dioctyl phthalate, dibutyl phtalate, phthalio acid benzyl butyl ester, and phthalic acid diisononyl ester Aliphatic-carboxylic-acid ester, such as dioctyl adipate, a succinic aoid screw (2-methylnonyl), dibutyl sebacate, and butyl oleate. Alcohol ester, such as pentaerythritol ester

,0055]Phosphoric ester, such as triootyl phosphate and tricresyl phosphate Epoxy plasticizers, such as epoxidized soybean oil, 4,5-epoxy hexahydrophthalic acid dioctyl, and epoxy stearic acid benzyl. Chlorinated paraffin. Polyester plasticizers, such as polyester dibasic acid and dihydric alcohol are

polyoxyalkylene, hydroxyl group content polyoxyalkylene may be sufficient and the polymer produced preferred. As such polyoxyalkylene, 1.5 or less polyoxyalkylene has preferred Mw/Mn which a with a eyanide complex etc. from the point that a low molecular weight body is hypoviscosity few. As such changed into the polymer which specifically closed terminal hydroxyl groups by hydrocarbon groups, molecular weights of 4000 or more thing was preferred, and was manufactured using the compound urethane bond, etc. is preferred. It is the polymer most preferably closed by the allyl group via the polystyrene, polybutadiene, alkyd resin, polychloroprene, polyisoprene, polybutene, hydrogenation by changing the hydroxyl group into other organic groups may be sufficient. Especially the thing such as an alkyl group and an alkenyl group, via combination of an ether bond, an ester bond, polyoxyalkylene polymer (A) and compatibility with (B) to polyoxyalkylene is [among these] [0056]Polyoxyalkylene, polyester, Polly alpha-methylstyrene, Polymerio plasticizers, such as polybutene, epoxidation polybutadiene, and Butadiene Acrylonitrile. The viewpoint of a

weatherability wants to improve, an adhesive property can be improved by considering it as the non-005/JThese plasticizers can be suitably chosen according to a use or the purpose. Although use of plastic combination with a large molecular weight uses what is called a polymeric plasticizer, and a plasticizer does not use a plasticizer rather rather than is necessarily required in the case of an high di-(2-ethylhexyl)phthalate of flexibility is the most common, For example, when high

mainly use plasticizers, such as an adhesives use, in particular, it is useful. The amount of the solvent used has 0.001 ~ 500 preferred weight section to a polyoxyalitylene polymer (A) and a total of 100 adjustment of viscosity, and the improvement in preservation stability of a constituent, and does not adjusting viscosity, in the non-plastic combination which can also add a solvent for the purpose of (Solvent) When using the constituent of this invention as a hardenability constituent again, when [0058]These plastioizers may be used independently and may use two or more sorts together. weight sections of (B).

ketone ester species, and ester ether can be used. When saving the constituent of this invention at a long period of time, since preservation stability of alcohols improves, they are preferred. As alcohols, alkyl alcohol of the carbon numbers 1-10 is preferred, and methanol, ethanol, isopropanol, isopentyl Alcohols, ketone, ester species, ether, ester alcohols, ketone alcohol, ether alcohol, ketone ether, [0059]As a solvent, aliphatic hydrocarbon, aromatic hydrocarbon, and halogenated hydrocarbon. alcohol, especially hexyl alcohol, etc. are preferred.

out 0.0001–10 weight-section use to a polyoxyalkylene polymer (A) and a total of 100 weight sections silicon group may be used. The following compound is mentioned as a concrete example. Those one hardening accelerator catalyst which promotes the hardening reaction of a hydrolytic basis content (Hardening accelerator catalyst) When stiffening the hardenability constituent in this invention, the sort or two sorts or more are used. As for a hardening accelerator catalyst, it is preferred to carry

[0061]As a Suzuki rate compound, dibuty/tin bisacety/lacetonate, dibuty/tin bis-ethylacetoacetate, a dibutyltin diacetate, dibutyltin mono- acetate, Organic tin carboxylate like dialkyl tin dicarboxylates, [0060]Specifically, the tin compound shown below is mentioned. Various compounds of divalent tin, Tetravalent tin various compounds, such as a reactant of dialkyl tinoxide, the reactant of an ester such as dibutyltin malate, or dialkyl tin mono- carboxylate, The Suzuki rate compounds, such as such as 2-ethylhexanoic acid tín, naphthenic acid tin, and stearic acid tín. Dibutyltin dilaurate, compound and dialkyl tinoxide, and an alkoxy silane compound, and a dialkyl tin dialkyl sulfide. dibutyltin monoacetyl acetonate monoalkoxide, etc. are mentioned. Carry out heating mixing, dialkyl tin bisacetylacetonate and a JIARUKIRUSUZU monoacetyl acetonate monoalkoxide,

etc. is also preferred, and can be used. The following are mentioned as a hardening accelerator which can be used besides a tin compound. Metal salt, such as an alkyl titanate, an organic silicon titanate, nade to react as a reactant of dialkyl tinoxide and an ester compound, and the tin compound made libutyltin oxide and phthalic ester, such as dioctyl phthalate and phthalic acid diisononyl ester, are liquefied is mentioned. In this case, as an ester compound, tetraethyl silicate, its partial hydrolysis [0062] The compound which reacted or mixed these tin compounds with low molecule alkoxysilane condensate, etc. can be used besides ester of aliphatic series and aromatic carboxylic acid organic-carboxylic-acid bismuth salt, and organic-carboxylic-acid lead salt.

chosphorio acid screw (2-ethylhexyl). A butylamine, hexylamine, octylamine, decyl amine, lauryl amine, [0063]Acidic compounds, such as phosphoric acid, p-toluenesulfonic acid, phthalic acid, and a

Hexamethylenediamine, diethylenetriamine, triethylenetetramine, Amine compounds, such as Aliphatic series monoamines, such as N,N-dimethyl- octylamine, ethylenediamine,

aminosilane coupling agents, such as aliphatic polyamine compounds, such as tetraethylenepentamine, an aromatic amine compound, alkanolamine, 3-(2-aminoethyl) amino-propyltrimethoxysilane, and 3-

temperature region comparatively especially. A hardening accelerator can be used combining one sort especially the 1st class amine compound, since their hardening facilitatory effect will improve, their concomitant use is preferred. By combining basic compounds, such as an above-mentioned acidic [0064]If a tin compound and a bismuth compound are used together with an amine compound, compound, an amine compound, etc., shows a higher hardening facilitatory effect in a high

(Storage stabilizer) A storage stabilizer can be added in order to improve the storage stability of the hardenability constituent of this invention further again. A storage stabilizer means the compound which can control hydrolysis of a polyoxyalkylene polymer (A) and (B) under existence of a small amount of water, or a compound with the high drying effect.

added at least, before using independently also being able to use these together and adding a curing polyoxyalkylene polymer (A) and a total of 100 weight sections of (B). A compound which generates weight section. Specifically, the following compounds are raised. It is preferred to be simultaneously strong acid by hydrolysis like mineral acid or chlorosilicane demonstrates the effect by one or less 0065]The amount of the storage stabilizer used has 0 - 30 preferred weight section to a

dimethyl malonate, and diethyl malonate etc. As mineral acid, it is mineral acid, such as chloride, sulfuric acid, and nitric acid, etc. [Me $_2$ CHO] $_2$ P(=O) (OH) [Me $_2$ CHO] P(=O) (OH) $_2$, (EtO) $_2$ P(=O) (OH) (EtO)P(=O) (OH) 2, (MeO) Phosphorio acid alkyl ester, such as 2P(=O) (OH) (MeO)P(=O) (OH) 2 and alcohol, n-amyl alcohol, Alcohols, such as isoamyl alcohol, 1-hexanol, octyl alcohol, 2-ethylhexanol, cellosolve, ethylene glycol, propylene glycol, glycerin, a diethylene glycol, and dipropylene glycol. [0066]Methanol, ethanol, 1-propanol, 2-propanol, 1-butanol, Isobutyl alcohol, 2-butanol, t-butyl acetoacetate, ethyl acetoacetate, and butyl acetoacetate, and an acetoacetic ester compound, [0067]Malonic ester compounds, such as 1,3-diketones, such as an acetylacetone, methyl catalyst

 $\{B_{LO}\}_2P(=0)$ (OH) (BuO)P(=0) (OH) $_2$ and $\{P_{LO}\}_2P(=0)$ (OH) (PrO)P(=0) (OH) $_2$ (In addition, a methyleur) group and Et express an ethyl group, Pr expresses a propyl group, Bu expresses a butyl group, and Me is the same as that of below.)

ORUTOGI acid alkyl ester, such as ORUTOGI acid trimethyl ** triethyl orthoformate, tripropyl orthoformate, and ORUTOGI acid tributyl

[0068]As altacetic acid alkyl ester, it is altacetic acid alkyl ester, such as altacetic acid trimethyl ** tetrapropylsilicate, and tetrabuthyl silicate. The partial hydrolysis condensate of tetramethyl silicate alt.acetic acid triethyl. Tetraalkyl silicate, such as tetramethyl silicate, tetraethyl silicate, and/or tetraethyl silicate.

trialkoxysilane, such as vinyltrimetoxysilane. Aryl trialkoxysilane, such as phenyltrimethoxysilane. Chlorosilicanes, such as methyltrichlorosilane, methyldi chlorosilicane, dimethyldichlorosilane, [0069]Alkyltrialkoxysilane, such as methyl trimetoxysilane and ethyltrimethoxysilane. Alkenyl

 CH_2 =Si(ON=CMe $_2$) $_2$, MeSi(NMe $_2$) $_3$, MeSi(ONMe $_2$) $_3$, MeSi(NMeC(=O) Me) $_3$, MeSi(OCMe=CH $_2$) $_3$. trimethylchlorosilane, phenyltrichlorosilane, and diphenyl dichlorosilane [0070]Silazanes, such as hexamethyldisilazane

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[0071] Hydrolytic organic titanium compounds, such as tetraisopropoxy titanium, tetrabutoxytitanium, and tetraoctyl oxytitanium, And those condensates, Ti(OCHMe₂) ₂[OCMe=CHCOMe] ₂, Ti(OBu) ₂ $[{
m Oc}_2{
m H_4}{
m N}({
m C}_2{
m H_4}{
m OH})_2^{}]_2$, Ti(OH) $_2^{}$ [OCHMeCOOH] $_2^{}$, Ti(OCHMe $_2^{}$) $_3^{}$ [OCOC $_1{
m PH}_{35}^{}$], Ti (OCHMe $_2^{}$)

 $[\mathsf{OCOC}_{\mathsf{I7}\mathsf{H}_{35}}]_3$, isopropoxy titanium dimethacrylate monoisostearate, isopropoxy titanium tris (4amino benzoate), isopropoxy titanium tris (dioctyl phosphate), etc.

[0072]Zeolite. Alkyl mercaptan, such as dodeoyl mercaptan and t-butyl mercaptan. Mercaptosilanes, such as gamma-mercapto propyltrimethoxysilane. Sulfhydryl group content carboxylic acid, such as

[0073]Suifhydryl group content ester compounds, such as 2-ethylhexyl thioglycolate. Sulfhydryl group content polymer like KAPUKYUA 3-800 (Diamond SHAMUROKKU Chemicals both-ends sulfhydryl group content polyoxyalkylene), a thiophenol, thiobenzoic acid, etc. 2-mercaptopropionic acid and thiosalicylic acid.

[0074]The partial hydrolysis condensate of the point of cost and an effect to vinyltrimetoxysilane, a tetraethoxysilane, tetramethyl silicate, and/or tetraethyl silicate is [among these] the most

Antioxidants generally used, such as each compound of a benzophenone series, a benzoate system, a using it especially combining two sorts or all from light stabilizer, an antioxidant, and an ultraviolet ray few effects of an improvement of weatherability, and if five weight sections are exceeded, there is no agent used, it is preferred that it is the range of 0.1 - 10 weight section to a polyoxyalkylene polymer and/or a phosphite system antioxidant are combined is effective. As for the amount of the antiaging (A) and a total of 100 weight sections of (B), respectively. In less than 0.1 weight sections, there are (Thixotropy grant agent) A thixotropy grant agent may be used again for the improvement of lappet systems, light stabilizer, and an ultraviolet ray absorbent are used suitably. These antiaging agents absorbent. Specifically, especially the thing for which the hindered amine light stabiliser of the 3rd method from being in the case of raising an effect as a whole taking advantage of each feature by nature. As such a thixotropy grant agent, hydrogenation castor oil, fatty acid amide, etc. are used. cyanoacrylate system, an acrylate system, a hindered phenol system, the Lynn system, and sulfur class or the 2nd class, a benzotriazol system ultraviolet ray absorbent, a hindered phenol system, have especially a preferred thing that it may use together and is used together. It is a desirable (Antiaging agent) As an antiaging agent again A hindered amine system, a benzotriazol system, great difference in an effect and it is economically disadvantageous.

NOKURAKKU SP, NOKURAKKU SP-N, NOKURAKKU NS-5, NOKURAKKU NS-6, NOKURAKKU NS-30, NOKURAKKU SO, NOKURAKKU NS-7, NOKURAKKU DAH (all are the products made from the Ouchi Shinko Chemical industry above), ADEKA stub AO-30, ADEKA stub AO-40, ADEKA stub AO-60, ADEKA stub AO-15, ADEKA stub AO-15, ADEKA stub AO-18, the ADEKA IRGANOX-1098, IRGANOX-1222, IRGANOX-1330, IRGANOX-1425WL (all are made in Tiba Specialty stub 328, ADEKA stub AO-37. (-- the above -- any ---) by Asahi Denka Kogyo K.K.. IRGANOX-245, IRGANOX-259, IRGANOX-565, IRGANOX-1010, IRGANOX-1035, IRGANOX-1076, IRGANOX-1081, preferred, and, specifically, the following oan be illustrated. NOKURAKKU 200, NOKURAKKU M-17, [0075]As an antioxidant, a hindered phenol system and/or a phosphite system anti-oxidant are Chemicals, Inc. above).

SANORU LS-292, SANORU LS-2626, SANORU LS-1114, SANORU LS-744 (all are the Sankyo Co., stub LA-601 (all are the Asahi Denka Kogyo K.K. make above), SANORU LS-770, SANORU LS-765, [0076]As light stabilizer, the 2nd class and/or the 3rd class hindered amine system compound have CHIMASSORB119floor line (all are made in Tiba Specialty Chemicals, Inc. above), ADEKA stub LA-57, ADEKA stub LA-62, ADEKA stub LA-67, ADEKA stub LA-63P, ADEKA stub LA-68LD, ADEKA them, and specifically, [preferred] Tinuvin 622LD, the tinuvin 144, CHIMASSORB944LD,

tinuvin 326, the tinuvin 327, the tinuvin 329, the tinuvin 213, and the tinuvin 120 (all are made in Tiba compound has it, and specifically, [preferred] The tinuvin P, the tinuvin 234, the tinuvin 320, the [0077]As an ultraviolet ray absorbent, a benzotriazol system compound or a benzoate system Ltd. make above), etc. can be illustrated.

curing-in-air compound, a photoresist compound) An air-oxidation hardenability compound and a Specialty Chemicals, Inc. above) can be illustrated. [0078]The tinuvin B5353 which is a mixture of the above-mentioned compound, the tinuvin B75 (all are made in Tiba Specialty Chemicals, Inc. above), etc. can be used.

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photoresist compound can be added in order to improve the adhesion of dust, and surface tackiness over a long period of time. Using together is more preferred although these compounds may be used independently. The amount used has 0.001 – 50 preferred weight section to a polyoxyalkylene polymer (A) and a total of 100 weight sections of (B).

oil and an isocyanate compound, the acrylic polymer by which conversion was carried out with drying conversion of the drying oil, and functionality polyoxyalkylene, The resultant (urethane oil) of drying polyester compound (air-drying nature unsaturated polyester) obtained by the polycondensation of Drying oil, such as tung oil, linseed oil, perilla oil, soybean oil, sunflower seed oil, and hempseed oil, oil, The epoxy resin by which conversion was carried out with drying oil, the silicon resin by which reaction by oxygen in the air as an air-oxidation hardenability compound is preferred. Specifically polybutadiene and diene of the carbon numbers 5–8, and a copolymer, the allyloxy group content mallein-ized denaturation, boiled oil denaturation, etc.) of this polymer or a copolymer, etc. are allyloxy group content glycol and polyvalent carboxylic acid --- the various denaturation things 0079]The compound which contains in intramolecular the unsaturation group which causes a The resultant of the various alkyd resins and drying oil which are produced by carrying out conversion was carried out with drying oil, Diene system polymers, such as a polymer of mentioned further.

Polyfunctional acrylate, such as a compound containing the acrylyl group produced by making a hydroxy compound, acrylic acid, and methacrylic acid, such as polyester polyol, react (meta), is used for a short time, and produces physical-properties change of hardening etc. by operation of light can [0080]As a photoresist compound, what molecular structure causes a chemical change considerably be used. Many things, such as a constituent which contains a monomer, oligomer, resin, or them as this kind of a compound, are known and marketed, and can use these publicly known compounds arbitrarily. Among these Polyhydric alcohol, such as trimethylolpropane, polyether polyol, the general-purpose.

reduction again. The effect of reducing a modulus by addition of these compounds without worsening (Modukus regulator) The compound which has one silanol group in intramolecular, or the compound which can generate the compound which has one slanol group in intramolecular can be added for stickiness of the surface is acquired. The amount used has 0-10 preferred weight section to a physical-properties adjustment of a hardened material and the purpose of surface stickiness polyoxyalkylene polymer (A) and a total of 100 weight sections of (B).

trimethylsilane, hexyloxy trimethylsilane, 2-ethylhexyloxy silane, n-octyloxy silane, 2-chloropropyl oxy alcohol, can be used for intramolecular. Specifically N-trimethylsilyl acetamide, hexamethyldisilazane, trimethylsilane, methoxy dimethyl phenylsilane, methoxy methyldi phenylsilane, phenoxydimethyl phenylsilane, etc. are mentioned. Ethylene glycol, propylene glycol, 1,3-propanediol, Dipropylene etherification of the hydroxyl group of multivalent hydroxy compounds, such as 1,6-hexanediol, compound which has one silanol group, trimethylsilyl ether, such as fatty alcohol and aromatic triphenylsilanol, etc. are mentioned to intramolecular. As a compound which can generate the trimethylsilane, Phenoxy trimethylsilane, 2-methylphenoxy trimethylsilane, 2-ohlorophenoxy glycol, 1,2-butanediol, 1,4-butanediol, The compound etc. which carried out trimethylsilyl Methoxy trimethylsilane and ethoxy trimethylsilane, isopropyloxy trimethylsilane, Butoxy [0081]As a compound which has one silanol group, a trimethyl silanol, triethyl silanol, a

glycerin, trimethylolpropane, pentaerythritol, and sorbitol, can be used. (in addition to this) Organic colors, such as inorganic pigments, such as iron oxide, chrome oxide, and retardant, can be used. The room-temperature-curing nature constituent of this invention is suitable for the use as which it can be used for sealant, a water blocking material, adhesives, a coating agent, etc., and the dynamic flattery nature to the sufficient cohesive force and adherend of especially the [0082]According to a use, publicly known additive agents, such as mildewproofing material and fire titanium oxide, and copper phthalocyanine blue, and Phthalocyanine Green, can be used as paints. hardened material itself is required. [Example] The example and comparative example of this invention are explained below. A part shows a weight shows the molecular weight converted from the hydroxyl value of the polyoxyalkylene polymer which has a hydroxyl group which is a raw material. Mw/Mn is the value measured by the gel weight section. In the example 1-1 to 1-6 of manufacture, a hydroxyl value conversion molecular

JP,2001-072855,A [DETAILED DESCRIPTION]

permeation chromatograph using a tetrahydrofuran as a solvent. The analytical curve was prepared using the correlation sample of styrene.

Next, the ally! chloride was made to react, the unreacted ally! chloride was removed and refined, and reactant, methyl dimethoxysilane which is a hydrosilyl compound was made to react under existence of a platinum catalyst, and the polypropylene oxide (P1) which has a methyl dimethoxy silyl group at the polypropylene oxide which has an allyl group at the end was obtained. it was alike, received this methanol solution of sodium methoxide was added, scale loss pressing-down methanol was distilled (Example 1-1 of manufacture) Use glycerin as an initiator and Under existence of a zinchexacyano off, and the terminal hydroxyl groups of polypropylene oxide was changed into sodium alcoholate. cobaltate glyme complex compound catalyst, To the hydroxyl value conversion molecular weight 17000 produced by making propylene oxide react, and polyoxypropylene triol of Mw/Mn=1.3. The the end was obtained.

ally! group at the end by the same method as the example 1 of manufacture was obtained using the zinchexacyano cobaltate glyme complex compound catalyst. The polypropylene oxide which has an made to react under existence of a platinum catalyst to this reactant, and the polypropylene oxide hydroxyl value conversion molecular weight 17000 produced by making propylene oxide react, and polyoxypropylene diol of Mw/Mn=1.3. Methyl dimethoxysilane which is a hydrosilyl compound was (Example 1-2 of manufacture) Use propylene glycol as an initiator and Under existence of a (P2) which has a methyl dimethoxy silyl group at the end was obtained.

under existence of a platínum catalyst, and the polypropylene oxide (P3) of the molecular weight 9000 continuously, it refined, and the polypropylene oxide (Mw/Mn=2.0) which has an allyloxy group at the potassium hydroxide catalyst, scale loss pressing-down methanol was distilled off to it, and terminal end was obtained. Methyl dimethoxysilane which is a hydrosilyl compound was made to react to this chlorobromomethane and performing polymers quantification, the allyl chloride was made to react polyoxypropylene diol of the hydroxyl value conversion molecular weight 3000 obtained using the (Example 1-3 of manufacture) The methanol solution of sodium methoxide was added to hydroxyl groups was changed into it at sodium alcoholate. Next, after making it react to which has a methyl dimethoxy silyl group at the end was obtained.

polymerization initiator, and the polypropylene oxide (P5) which has a trimethoxysilyl group at the end polyoxypropylene diol of Mw/Mn=1.3. To this reactant, 3-mercapto propyltrimethoxysilane which is a cobaltate glyme complex compound catalyst. The polypropylene oxide which has an allyl group at the conversion molecular weight 17000 produced by making propylene oxide react, and polyoxypropylene was made to react using the 2,2'-azobis 2-methylbutyronitrile which is a polymerization initiator, and allyl group at the end by the same method as the example 1 of manufacture was obtained using the zinchexacyano cobaltate glyme complex compound catalyst, The polypropylene oxide which has an triol of Mw/Mn=1.3. To this reactant, 3-mercapto propyltrimethoxysilane which is a silyl compound (Example 1-4 of manufacture) Use glycerin as an initiator and Under existence of a zinchexacyano hydroxyl value conversion molecular weight 17000 produced by making propylene oxide react, and end by the same method as the example 1 of manufacture was obtained using the hydroxyl value (Example 1-5 of manufacture) Use propylene glycol as an initiator and Under existence of a silyl compound was made to react using the 2,2'-azobis 2-methylbutyronitrile which is a the polypropylene oxide (P4) which has a trimethoxysilyl group at the end was obtained.

polyoxypropylene triol of Mw/Mn≕1.2. gamma−isocyanate propyltrimethoxysilane was added to this, was obtained. (Example 1-6 of manufacture) It refined, after having used glyoerin as the initiator, performing the polymerization of propylene oxide under existence of a zinchexacyano cobaltate glyme complex compound catalyst and obtaining the hydroxyl value conversion molecular weight 10000 and the urethane-ized reaction was performed, and the polypropylene oxide (P6) which has a trimethoxysilyl group at the end was obtained.

of a zinchexacyano cobaltate glyme complex compound catalyst, The polypropylene oxide (G1) which (Example 2-1 of manufacture) Use polymeric plasticizer glycerin as an initiator and Under existence has an allyl group at the end by the same method as the example 1 of manufacture was obtained using the hydroxyl value conversion molecular weight 17000 produced by making propylene oxide react, and polyoxypropylene triol of Mw/Mn=1.3.

(Example 2-2 of manufacture) It refined, after having used polymeric plasticizer glycerin as the

nitiator, performing the polymerization of propylene oxide under existence of a zinchexacyano cobaltate glyme complex compound catalyst and obtaining the hydroxyl value conversion molecular weight 10000 and polyoxypropylene triol of Mw/Mn=1.2. (G2)

(Example 3–1 of manufacture) Add 0.2 mol of dibutyltin oxide to toluene 150cm³ in polymer hardening accelerator catalyst glass reactors, and 0.1 mol of 2-ethylhexanol is added, It was made to react removing the water which carries out azeotropy to bottom toluene of heating chuming until the water removing the emount of theories finishes distilling off. Then, 0.1 mol of acetylacetones are added, and it was made to react removing the water which carries out azeotropy to toluene further until the water of the amount of theories finishes distilling off. It filtered in order to remove a little sediments, and it distilled off under decompression of toluene further, and the light yellow tin compound (J1) was obtained.

trimethylsilyl group with a nuclear magnetic resonance spectrum, After the saturation sodium chloride mostly until it filtered the reaction mixture, the water layer checked filtrate with the pH test paper in agitator, and 3.3 mol of sodium hydroxide ground further was added, 3.0 mol of trimethylchlorosilanes sodium sulfate was put into the organic layer, and it dried to it. Filtration removed anhydrous sodium aqueous solution washed after washing enough after checking that the reaction had been completed hardening accelerator catalyst flowing-back condenser tube and an agitator. 1 mol of dibutyltin oxide sulfate, distillation refining was carried out under decompression, and the trimethylsilyl ether (L1) of emperature up was carried out to 40 ** after the end of dropping, and it agitated for 8 hours. Fine carried out to epoxy curing agent glass reactors, The pyrogenetic reaction was carried out at 90 ** order of ammonium chloride solution and ion exchange water and it became pH7 mostly, anhydrous and 0.5 mol of 2-ethylhexanoic acid ethyl were added, and it heated at 120 ** for 5 hours, agitating (Example 5-1 of manufacture) 1 mol of trimethylolpropane was put into the glass reaction vessels Example 4-1 of manufacture) Weighing of epicure H-1(epoxy resin hardener by oil recovery shell spoxy company)2 mol and the KBM403 (epoxysilane by Shin-etsu chemicals company) 2.5 mol is which attached the modulus regulator dropping funnel, the flowing-back condenser tube, and the were dropped over 1 hour from the dropping funnel at the bottom room temperature of churning. Example 3-2 of manufacture) In the 3 Thu mouth flask of the glass furnished with a polymer under a nitrogen atmosphere, and the tin compound (J2) of uniform light yellow was obtained. sodium chloride has deposited in a reaction vessel as a reaction progresses. By analyzing a for 5 hours, agitating under a nitrogen atmosphere, and the compound (K1) was obtained. trimethylolpropane was obtained.

(Example 5–2 of manufacture) To the glass reaction vessels which attached the modulus regulator dropping funnel, the flowing–back condenser tube, and the agitator. 1.5 mol of 2–ethylhexyl alcohol was put in, and 0.5 mol of trimethylohlorosilane and the mixture of 0.5 mol of hexamethyldisilazane were dropped over 2 hours under churning at the room temperature. It agitated at 40 ** after the end of dropping for 5 hours, 0.2 more mol of methanol was added, and it allowed to stand at the room temperature all night. After it removed the ammonium chloride which carried out the byproduction by filtration on the next day, and ammonium chloride solution washed filtrate and drying and filtering with anhydrous sodium sulfate, distillation refining was carried out under decompression and the trimethylsilyl ether (L2) of 2–ethylhexyl alcohol was obtained.

(Example 1 of an examination) Polypropylene oxide (P1) – (P6) manufactured in the example 1–1 to 1–6 of manufacture and an antiaging agent, and a thixotropy grant agent to what carried out mixture dispersion. The bulking agent and paints from which moisture was beforehand removed by stoving were added, and it mixed, and Epicoat 828 (bisphenol A type epoxy resin by an oil recovery shell epoxy company), the epoxy curing agent, the plasticizer, and the storage stabilizer were added further, and it mixed. Furthermore the silane compound and the curing catalyst were added, mixing and after defoaming, the cartridge for sealing materials which can interrupt penetration of the hygroscopic surface moisture in the air was filled up, and the constituents D1–D8 of this invention and the comparative constituents D9–D12 were obtained. The kind and quantity of each raw material are as having been shown in Table 1.

are as navitable point and the ease (handling nature) of dealing with it of a class product was contrible out (O which is especially easy to deal with it, :O which is easy to deal with it, and which it, and which it, and which has the low viscosity of a constituent is dealt with generally, and a sex is good). A result is shown in Table 1-1.

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[0085] [Table 1] ï

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[0086]The following examinations were done about the constituent (11 points) which was good among the obtained constituents as for handling nature. A result is shown in Table 1–2.

(Tuck free examination) The tuck free examination was done based on JIS A5758. The class product was extruded from the cartridge, and promptly, on the glass plate, the spatula was used for about 3 mm in thickness, and it accustomed level, and placed at the normal condition. The time which it took that a constituent stops adhering to a fingertip since the surface was lightly touched and accustomed it by the fingertip purified by ethanol level was measured, and it was considered as tack free time. (Tension shear test) In [extrude a class product from a cartridge and] the bottom of 50% of humidity at 23 **. To an aluminum plate (25 mm in width, 100 mm in length, and 3 mm in thickness, and adhesion area of 25 mm in 25 mm, it applied so that it might be set to about 0.2 mm in thickness, and in 30 seconds, one more piece of aluminum plate is piled up alternately, and was stuck, and tensile shear bond strength (unit kg/cm²) was measured after care of health for seven days on the fone?

[Table 2]

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added, and it mixed, and the plasticizer, the storage stabilizer, and the epoxy curing agent were added was made into A agent. To what added the polypropylene oxide (P1) manufactured in the example 1-[0088](Example 2 of an examination) A plasticizer, a bulking agent, and paints were added to Epicoat further. The bulking agent and paints from which moisture was beforehand removed by stoving were further, and it mixed. Furthermore the silane compound and the curing catalyst were added, mixing and defoaming of were done, and this was made into B agent. The kind and quantity of each raw I to 1-6 of manufacture - (P6) the modulus regulator, and the curing-in-air compound, was mixed, 828 (bisphenol A type epoxy resin by an oil recovery shell epoxy company), it mixed to it, and this and added and carried out mixture dispersion of an antiaging agent and the thixotropy grant agent material are as having been shown in Table 2.

out (O which is especially easy to deal with it, :O which is easy to deal with it, :x which is hard to deal three-stage evaluation of the ease (handling nature) of dealing with it of a class product was carried with it, and * — the thing which has the low viscosity of a constituent is dealt with generally, and a [0089]A agent and B agent which were obtained were mixed and agitated just before use, and the constituents D13-D20 of this invention and the comparative constituent D21 $^-$ 24 were obtained sex is good). A result is shown in Table 2-1. http://www4.ipdl.inpit.go.jp/cgi-bin/tran_web_cgi_ejje?atw_u=http%3A%2F%2Fw2Fw4.ipdl.i.. 2010/03/03

[0600]

JP,2001-072855,A [DETAILED DESCRIPTION]

Table 3

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[0091]The following examinations were done about the constituent (11 points) which was good among the obtained constituents as for handling nature. A result is shown in Table 2-2.

and accustomed it by the fingertip purified by ethanol level was measured, and it was considered as which it took that a constituent stops adhering to a fingertip since the surface was lightly touched mixing, on the glass plate, the spatula was used for about 3 mm in thickness, and base resin and a (Tuck free examination) The tuck free examination was done based on JIS A5758. Uniformly, after hardening agent were sounded level, and were placed promptly at the normal condition. The time tack free time.

(Tension shear test) In [mix A agent and B agent uniformly, and] the bottom of 50% of humidity at

2010/03/03

cond strength (unit kg/cm^2) was measured after care of health for seven days on the conditions with 23 **, To an aluminum plate (25 mm in width, 100 mm in length, and 3 mm in thickness), an adhesion area of 25 mm x 25 mm, it applied so that it might be set to about 0.2 mm in thickness, and in 30 seconds, one more piece of aluminum plate is piled up alternately, and was stuck, and tensile shear 5-mm the speed of testing for /.

[0092]

Table 4

1

22 D22 D23 88 22 88 28 2 88 ទូន 77.5 5 3 018 D19 22 27 75 37.5 37.5 37.5 88 -1 -1 1.0 0.1 0.7 0.8 35 31 38 38 7.5 88 88 유학 = 7 88 Ξ 7## 2 7 1452 '18 7## 2 7 7453 '19 4 # 2 7 720 '20 7 # # 1 * 7 7 7 1010 'El テンクリン・ナイ (5r) 引送記載機器(3E) (4g/cm) *华心多野 \$5. 50\$50 park +# 11:1252171 70-5 28009 29 チャンインジューンティング・ファンド TEST - CAT 27.7.4.10 01.4.7.4.10 02.4.2.4.10 81. SST 11. SST **7**3 2020 88 뾽 エポキシ出版(C) 充拠別 变死硬化性化合物 无键化性化合物 **ソラン広会協(?)** 年がよう 4間報金 女女な様にサイ **伊莱安斯斯** 老化协业和 翼 平 五 **光雕**型 KREA

[0093]

[Effect of the Invention]As explained above, according to this invention, the polymer which has a hydrolytic silicon group industrially obtained easily considering the silane compound [handling / a silane compound] as a raw material is used, It is dealt with by hypoviscosity, and excels in a sex, and

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JP,2001-072855,A [DETAILED DESCRIPTION]

with adherend, and, moreover, the hardenability constituent which can be adjusted in the range with a wide cure rate can be obtained. excels in the mechanical strength of the hardened material after hardening, and adhesive strength

[Translation done.]

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